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Peterson J., DeBoy R., Dodson R., Gwinn M., Haft D., Hickey B., Kolonay J.F., Nelson W.C., Umayam L.A., Ermolaeva M., Salzberg S.L., Delcher A., Utterback T., Weidman J., Khouri H., Gill J., Mikula A.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Whole genome comparison of Mycobacterium tuberculosis clinical and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MEDLINE-9829599; PubMed-9634230;
Cole S.T., Brosch R., Parkhill J., Garnier T., Churcher C., Harris Gordon S.T., Brosch R., Parkhill J., Garnier T., Churcher C., Harris Gordon S.V., Eighmeier K., Gas S., Barry C.E. III, Tekala F., Badcock K., Basham D., Brown D., Chillingworth T., Connor R., Davies R., Davies R., Broyln K., Feltwell T., Gentles S., Haminn N., Holroyd Hernsby T., Jagels K., Krogh A., Motlean J., Moule S., Murphy L., Oliver S., Seeger K., Skelton S., Rajandram M.A., Rogers J., Rutter S., Seeger K., Skelton S., Squares S., Squares R., Sulston J.E., Taylor K., Whitehead S., Barrell B.G.;
Pocipheiring the biology of Mycobacterium tuberculosis from the Complete genome sequence."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Actinomycetales;
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Submitted (APR-2001) to the EMBL/GenBank/DDBJ databases.
-!- SIMILATY: STRONG, TO M.SMEGMATIS MIHF.
EMBL; 280108; CAB02193.1; --
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TIGR; WILL431; -.
TUGR; WILL431; -.
Tubercutist; Rv1388; -.
Hypothetical protein; Complete protecome.
Hypothetical protein; Complete protecome.
Hypothetical 190 AA; 20835 MW; 376672DCF96AF60D CRC64;
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01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-OT-2000 (TrEMBLrel. 13, Last annotation update)
Hypothetical protein SCO1504.
SCO1504 OR SC9C5.28.
Streptomyces coelicolor.
                                                                                                                                                                                                                                                                                                                                          P71658 PRELIMINARY; PRT; 190 AA. P71658; P71658; P71658; P71678; P7167
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Corynebacterineae; Mycobacteriaceae; Mycobacterium
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Mismatches
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EMBL; AE007015; AAK45698.1; ALT_INIT.
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STRAIN-CDC 1551 / Oshkosh;
Fleischmann R.D., Alland D
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   9; Conservative
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                                                                           7 AAARRARAE 15
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Best Local Similarity
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Q9KXP6;
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                                           Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Pedulla M.L., Lee M.H., Lever D.C., Hatfull G.F.;
"A novel host factor for integration of mycobacteriophage L5.";
Submitted (AGG-1998) to the EMBL/GenBank/DDBJ databases.
EMBL; U75344; AAC2846.1; --
SEQUENCE 105 AA; 11635 MW; B73846DBFA6CA838 CRC64;
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Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Corynebacterineae; Mycobacteriaceae; Mycobacterium.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Mycobacterium smegmatis.
Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Corynebacterineae; Mycobacteriaceae; Mycobacterium.
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1-OCT-2001 (TrEMBLrel. 18, Last annotation update)
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Last annotation update)
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Nature 409:1007-1011(2001).
EMBL; AL583918; CAC30048.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          56.2%; Score 9; DB 16; 100.0%; Pred. No. 2.9;
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100.0%; Pred. No. 2.9
tive 0; Mismatches
       Pred. No.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Created)
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       Best Local Similarity 100.0%;
Matches 10; Conservative 0
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Best Local Similarity 100...
Second 9; Conservative
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                                                                                                                       1 ARRAARAAR 10
                                                                                                                                                                                               71 ARRAARAAR 80
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Barrell B.G.;
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20 RRAARAA 27
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09x696;
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Q9X696
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                                                                                                                                                                                                                                                                                                                                                                                                                 Redenbach M., Kieser H.M., Denapaite D., Eichner A., Cullum J., Kinashi H., Hopwood D.M.;
"A set of ordered cosmids and a detailed genetic and physical map for the 8 Mb Streptomyes coeliclor A3(2) chromosome.";
Mol. Microbiol. 21:77-96(1996).
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MEDLINE-21996410; PubMed-12000953;
MEDLINE-21996410; PubMed-12000953;
Bentley S.D., Chatler K.F., Cardeno-Tarraga A.-M., Challis G.L.,
Thomson N.R., James K.D., Harris D.E., Quail M.A., Kieser H.,
Harper D., Bateman A., Brown S., Chandra G., Chen C.W., Collins M.,
Cronin A., Fraser A., Goble A., Hidalgo J., Hornsby T., Howarth S.,
Huang C.-H., Kieser T., Larke L., Murphy L., Ollver K., O'Neil S.,
Rabbinowitsch E., Rajandream M.A., Rutherford K., Rutter S.,
Seeger K., Saunders D., Sharp S., Squares R., Squares S., Taylor K.,
Watren T., Wietzorrek A., Woodward J., Barrell B.G., Parkhill J.,
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Sukryyota Metazoa; Chordata; Cranlata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Musinae; Mus
NCBL_raxID-10090;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Complete genome sequence of the model actinomycete Streptomyces
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Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Streptomycineae; Streptomycetaceae; Streptomyces.
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                                                                                                                                                                                                                                                                   Cerdeno A.M., Parkhill J., Barrell B.G., Rajandream M.A.; Submitted (JUN-2000) to the EMBL/GenBank/DDBJ databases.
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01-MAR-2003 (TTEMBLTel. 23, Last annotation update)
Hypothetical type I antifreeze protein containing protein
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                                                                                                                                              Brown S.P., Harris D.;
Submitted (JUN-2000) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11 protein; Complete protecme.
750 AA; 81839 MW; 469E5F417EDC8862 CRC64;
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tive 0; Mismatches
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EMB., ALS9109; CAB93184.1. -.
InterPro: IPRO00767; Disease_resist.
PRINTS; PR00364; DISEASERSIST.
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STRAIN-C57BL/6J; TISSUE-EYe;
MEDLINE-22334683; PubMed-12466851;
                                                                                                                                                                                                                                                                                                                                                                   STRAIN-A3(2);
MEDLINE-97000351; PubMed-8843436;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match 56.2
Best Local Similarity 100.
Matches 9; Conservative
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76 AARAAARRA 84
                                                                                             SEQUENCE FROM N.A.
                                                                                                                                                                                                                  SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE FROM N.A.
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                                                WCBI_TaxID-1902
                                                                                                                       STRAIN-A3(2);
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01-Mar-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (Fragment)
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"Oryza sativa (japonica cultivar-group) genomic DNA, chromosome 1, PAC
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01-UUN-2002 (TrEMBLrel. 21, Last sequence update)
01-OCT-2002 (TrEMBLrel. 22, Last annotation update)
01-OCT-2002 (TrEMBLrel. 22, Last annotation update)
00684C02.19 protein.
07yza sativa (japonica cultivar-group).
Eukaryote; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnollophyta; Liliopsida; Poales; Poaceae; Enhartoldeae; Oryzeae; Oryza.
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Bacteria; Actinobacteridae; Actinomycetales; Streptomycineae; Streptomyces.
                                                                                                                                                                           Length 101;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 10; Length 167;
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EMBL; AP003290; BAB89063.1; -.
Gramene; Q8S1J9; -.
SEQUENCE 167 AA; 18542 WW; OC40040DFB49021B CRC64;
                                                                                                                       CBDCDC2D530ABB15 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   01-NOV-1999 (TrEMBLrel. 12, Created)
01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
                                                                                                                                                                      DB 11;
                                                                                                                                                                                                                                                                                                                                                                                                                                    167 AA.
                                                                                                                                                                      50.0%; Score 8; DB 1:
100.0%; Pred. No. 16;
1ve 0; Mismatches
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tive 0; Mismatches
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"Characterization of a vanillic acid non-
gene cluster from Streptomyces sp. D7.";
Microbiology 145.2393-2403(1999).
EMBL; AF134589; AAD28781.1;
InterPro; IPR003382; Flavoprotein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             STRAIN-cv. Nipponbare;
Sasaki T., Matsumoto T., Yamamoto K.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         PRT;
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60,770 full-length cDNAs.";
Nature 420:563-573(2002).
EMBL; AK021392; BAC25654.1; -.
Hypothetical protein.
NON_TER
                                                                                                                       SEQUENCE 101 AA; 10110 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             50.0%;
                                                                                                                                         Query Match
Best Local Similarity 100...
Best Conservative
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Best Local Similarity 100.
Matches 8; Conservative
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MEDLINE-99235742; PubMed-10217753;
MEDLINE-99235742; Mohn W.W.;
Martin V.J., Mohn W.W.;
Martin V.J., Mohn W.W.;
An ovel aromatic-ring-hydroxylating dioxygenase from the diterpenoid-
a novel aromatic-ring-hydroxylating dioxygenase from the diterpenoid-
degrading bacterium Pseudomonas abietaniphila BKME-9.";
J. Bacteriol. 181:2675-2682(1999).
EMED: ANDIOS1; ANDIOS2; FAA_hydrolase.
Interpro: IPR00253; FAA_hydrolase;
STOPENOIS57; FAA_hydrolase;
Pfam; PF01557; FAA_hydrolase;
SEQUENCE 336 AA; 37494 MW; CD221163FB1B44FB CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 STRAIN-BKME-9;
STRAIN-BKME-9;
Mohn W.W., Wilson A.E., Bicho P., Moore E.R.B.;
Mohya W.W., Wilson A.E., Bicho P., Moore E.R.B.;
Physiological and Phylogenetic Diversity of Bacteria Growing on Resin Acids.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Mus musculus (Mouse).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostoml;
Mammalia; Eutheria; Rodentia; Sciurognathl; Muridae; Murinae; Mus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Pseudomonas abietaniphila.
Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales;
Pseudomonadaceae; Pseudomonas.
NCBI_TaxID=89065;
DB 10; Length 274;
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01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-NAR-2003 (TrEMBLrel. 23, Last annotation update)
MHC class I H2-K gene (Haplotype d) (Fragment).
                                                                                                                                                                                                                                                                                                                                                                                  50.0%; Score 8; DB 2; 100.0%; Pred. No. 43;
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0; Mismatches
                                                                                                                                                                                                                                                                                                                                                     336 AA
         50.0%; Score 8; DB 10
100.0%; Pred. No. 37;
                                                                0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Syst. Appl. Microbiol. 0:0-0(1999).
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Best Local Similarity luv...
8; Conservative
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                                                                                Conservative
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81 AAARRARA 88
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                   Query Match
Best Local Similarity
Matches 8; Conserv
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09X4X0;
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Q31191
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Q9X4X0
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Okamoto O.K., Hastings J.W.;
"Circadian oscillations in the transcriptome of dinoflagellate cells: towards the clock circuitry.;
Submitted (APR-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AF508261; AA014680.1;
SEQUENCE 274 AA; 28168 MW; B5F522A5D45AC8BA CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
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                                                                                                                                                                                                                                                                     Gaps
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NCBI_TaxID=2972;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB 12; Length 204;
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EMBL. 13173; AACS4557.1; -.
InterPro; IPR005039; Sug_transporter.
InterPro; IPR005839; Sug_transporter.
Pfam; PF03169; Herpes_UL3; 1.
PROSITE; PS00216; SUGAR_FRANSPORT_1; 1.
SEQUENCE 204 AA; 21792 MW; 9292E6A8AA2CB8C6 CRC64;
                                                                                                                                                                                                                                                                     Indels
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01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
Chlorophyll A-C binding protein.
Pyrocystis lunula.
                                                InterPro; IPR004507; Ubix.
Pfam; PF02441; Flavoproteán; 1.
TIGRFAMB; TIGR00421; Ubix; 1.
SEQUENCE 200 AA; 21660 MW; 83D1B5C22C13CD3E CRC64;
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Last annotation update)
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                                                                                                                                                                                                       Query Match 50.0%; Score 8; DB 2; Best Local Similarity 100.0%; Pred. No. 28; Matches 8; Conservative 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          0; Mismatches
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Pred. No.
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100.0%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       bovine herpesvirus-1.";
Virology 213:28-37(1995).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Local Similarity 100.
les 8; Conservative
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                                                                                                                                                                                                                                                                                                                                               3 RAARAAAR 10
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Length 336; Indels

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Oryza sativa (japonica cultivar-group).
Wakaryota, Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoideae; Oryzeae; Oryza.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SEQUENCE FROM N.A.
STRAIN-16M / ATCC 23456 / Blotype 1;
MEDLINE-20020109; Pubmed-11756688;
Delyecchio V.G., Kapatral V., Redkar R.J., Patra G., Mujer C., Ivanova N., Anderson I., Bhattacharyya A., Lykidis A., Reznik C Jablonski L., Larsen N., D'Souza M., Bernal A., Mazur M., Golte Selkov E., Elzer P.H., Hagius S., O'Callaghan D., Letesson J.-. The genome sequence of the facultative intracellular pathogen Brucella melitensis.";
                                                                                                                                                                                                                                                                                                                                      SEQUENCE FROM N.A.
STRAIN-cv. Nipponbare;
Sasaki T., Matsumoto T., Yamamoto K.;
Cyza astiwa nipponbare(GA3) genomic DNA, chromosome 1, PAC
clone:P0699H05.
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Brucellaceae; Brucella.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   50.0%; Score 8; DB 10; Length 451; 100.0%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 710;
                                                                                                                                                                                                                                                                                                                                                                                                                                                   Submitted (FEB-2001) to the EMBL/GenBank/DDBJ databases. EMBL; - AP003299; BABB9880.1; - Gramene: Q85112; - SEQUENCE 451 AA; 49641 MW; 2E196EAA1DCB1468 CPCF64.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Indela
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          2E196EAA1DCB1468 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       710 AA; 76150 MW; 5DCA5720C36DBBFA CRC64;
                                                                                           01-07N-2002 (TrEMBLrel. 21, Created)
01-07N-2002 (TrEMBLrel. 21, Last sequence update)
01-0CT-2002 (TrEMBLrel. 22, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    01-WAR-2002 (TrEMBLrel. 20, Last sequence update) 01-UUN-2002 (TrEMBLrel. 21, Last annotation update) ATP-dependent helicase HRPB.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 U.S.A. 99:443-448(2002).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     50.0%; Score 8; DB 16;
100.0%; Pred. No. 79;
tive 0; Mismatches
                                                                  451 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    710 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Pred. No. 55;
0; Mismatches
                                                           PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Created)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 InterPro; IPR001410; DEAD.
InterPro; IPR001650; Hellcase_C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          EMBL; AE009614; AAL52999.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity 100.00
                                                        PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity 100.
Matches 8; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Pfam; PF00271; helicase_
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              237 RRAARAA 244
                                                                                                                                                           P0699H05.2 protein.
P0699H05.2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     2 RRAARAAA 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Brucella melitensis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Complete proteome.
SEQUENCE 710 AA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NCBI_TaxID-29459;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    01-MAR-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Q8YER0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RESULT 15
Q8YER0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         STRAIN=ATCC 15692 / PAO1;
BDLINE-20437337; PubMed=10984043;
Stover C.K., Pham X.-Q.T., Ervin A.L., Mizoguchi S.D., Warrener P. Hickey M.J., Brinkman F.S.L., Bufnagle W.O., Kowalik D.J., Lagrou Garber R.L., Goltry L., Tolentino E., Westbrock Wadman S., Yuan Y.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales;
-i- SUBUNIT: DIMER OF ALPHA CHAIN AND A BETA CHAIN (BETA-2-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    S., Wu Z., Paulger K., Lin
S., Wu Z., Paulsen I.T.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Garber R.L., Goltry L., Tolentino E., Westbrock-Wadman S., Brody L.L., Coulter S.N., Folger K.R., Kas A., Larbig K., Smith K.A., Spencer D.H., Wong G.K.-S., Wu Z., Paulsen I. Reizer J., Saler M.H., Hancock R.E.W., Lory S., Olson M.V. "Complete genome sequence of Pseudomonas acruginosa PAOI,
                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 356
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 16; Length 383; . 48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      le W.O., Kowalik D.J
, Westbrock-Wadman S
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Indels
                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE 356 AA; 39888 MW; 7BFB4957212E3F2D CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   383 AA; 42550 MW; 8DE3B490DD83C9B1 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           01-MAR-2001 (TrEMBLrel. 16, Created)
01-MAR-2001 (TrEMBLrel. 16, Last sequence update)
01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
Probable multidrug resistance efflux pump.
                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 7;
                                                                                                                                                                                                                                                                                                                                                                                                                                          50.0%; Score 8; DB 7
100.0%; Pred. No. 45;
vative 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   50.0%; Score 8; F
100.0%; Pred. No.
                                  (BY SIMILARITY)
                                                                                                        InterPro; IPR007110; Ig-11ke.
InterPro; IPR003597; Ig_c1.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR001039; MHC_I.
                                           EMBL; K01182; AAA39653.1; --
                                                                                                                                                                                                                                                                                                        PROSITE; PS50835; IG LIKE; 1
PROSITE; PS00290; IG MHC; 1.
Glycoprotein; Transmembrane.
                                                                                                                                                                                                                                          PRINTS; PR01638; MHCCLASSI.
ProDom; PD000050; MHC_I; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Pfam; PF00529; HlyD; 1.
PRINTS; PR01490; RIXTOXIND.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity 100.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      interPro; IPR006143; HlyD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Ouery Match
Best Local Similarity 100.
Matches 8; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          opportunistic pathogen."
Nature 406:959-964(2000)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Beudomonas aeruginosa
                                                                                                                                                                                                                                                                                      SMART; SM00407; IGC1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                3 RAARAAAR 10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              20 RARRAAR 27
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          187 ARAAARRA 194
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5 ARAAARRA 12
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Complete proteome.
SEQUENCE 383 AA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NCBI_TaxID-287;
                                                                                                                                                                                                 Pfam; PF00047
                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
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RESULT 13 Q914A1 ID Q914A

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, Goltsman E.,

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RESULT 14

Reznik G.

156 AARAAARR 163

Search completed: August 9, 2003, 16:32:58 Job time: 41.4571 secs